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# (54) Isolation and sequencing of the hazel FAd2-N gene

(57) The invention relates to the isolation from hazel (Corylus avellana L.) of the FAD2-N gene coding for the Δ12 desaturase enzyme of the microsomal fraction and, in particular, provides the nucleotide sequence and the deduced amino-acid sequence of the gene and provides for its use as a probe for the isolation of other plant desaturases. It also relates to the use of this gene for altering the desaturase levels and consequently the fatty-acid composition of the plant.

#### Description

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The present invention relates to the isolation from hazel ( $Corylus\ avellana\ L$ .) of the FAD2-N gene which codes for the  $\Delta$ 12 desaturase enzyme of the microsomal fraction.

More particularly, the invention relates to the nucleotide sequence, to the derived amino-acid sequence of the gene, and to its use as a probe for the isolation of other plant desaturases. It also relates to the use of this gene for altering the desaturase levels, and consequently the fatty-acid composition of the plant.

Alteration of the fatty-acid composition may have various applications in the industrial field. One of the greatest problems with hazelnuts is that they become rancid by oxidation. This is due to the auto-oxidation of unsaturated lipids with the consequent formation of volatile substances with a rancid odour which cannot easily be eliminated by the usual preservation systems. Amongst the possible strategies for reducing the tendency to become rancid, the best seems to be that of reducing the degree of unsaturation of the fatty acids present in the kernel oil, since susceptibility to auto-oxidation is positively correlated with this parameter. In fact, the rate of peroxide formation is correlated with the number of C=C double bonds in the fatty acids. The rate of auto-oxidation of the fatty acids in comparison with the oleate (18:1) is about 30 times greater in the linoleate (18:2) and 80 times greater in the linolenate (18:3). Moreover, the volatile substances resulting from the degradation of the linoleate and of the linolenate have a lower threshold of perception than those derived from the oleate. A reduction in linoleic acid should reduce the availability of substrates for lipoxygenase, reduce the loss of vitamin E during preservation, and reduce the production of volatile substances such as hexanals.

In the angiosperms, most of the synthesis of polyunsaturated lipids takes place by means of a single enzyme, that is,  $\Delta 12$  (or  $\omega 6$ ) desaturase (18:1 desaturase), of the endoplasmic reticulum, although there is an 18:1 chloroplast desaturase in the leaves of some plants. Moreover, this enzyme is responsible for more than 90% of the synthesis of polyunsaturated fatty acids in non-photosynthetic tissues such as, for example, in the kernels. The conversion of oleic acid (18:1) to linoleic acid (18:2) thus takes place by means of  $\Delta 12$  desaturase, and from linoleic acid to linolenic acid (18:3) by means of  $\Delta 15$  (or  $\omega 3$ ) desaturase.

It has been shown with mutants of *Arabidopsis* that the FAD2 locus contains a gene which codes for the oleate desaturase enzyme of the endoplasmic reticulum (Okuley et al, 1994, The Plant Cell 6, 147-158). The FAD2 gene was in fact able to complement mutants of *Acabidopsis* which were deficient in desaturase activity of the endoplasmic reticulum. The gene coding for the same enzyme in soya has also recently been isolated and sequenced (Heppard et al, 1995, Plant Physiol., in press).

A reduction in the  $\Delta 12$  desaturase levels should therefore lead to a reduction in the linoleic acid content and, as a secondary effect, probably also to a reduction in linolenic acid. In hazelnuts the percentage of linoleic acid varies from 5 to 15%; the percentage of linolenic acid is from 0.1 to 0.2%. A reduction in these fatty acids should therefore be useful in the preservation of hazelnuts. There is therefore clearly a need to isolate the gene which codes for the  $\Delta 12$  desaturase of the endoplasmic reticulum. The sequence of the gene could thus be used for gene inactivation in hazelnut kernels. This inactivation could be carried out either by the antisense technique (Smith et al. (1988) Nature 334, 724-726) or by the "transwitch" technique (Flavell (1994) Proc. Natl. Acad. Sci. USA 91, 3490-3496). In the antisense technique, the hazel would have to be transformed by the entire FAD2-N gene or by portions thereof, inserted in the opposite direction to the regulating sequences. In the "transwitch" technique, the hazel would have to be transformed by an identical copy of the FAD2-N gene.

The subjects of the present invention are defined by the following claims.

Embodiments of the present invention will now be described with reference to the following drawings, in which:

Figure 1 shows the restriction map of the N2 genome clone,

Figure 2 shows the nucleotide sequence of the hazel FAD2-N gene; the amino-acid sequence of the coding portion is also shown;

Figure 3 shows the nucleotide sequence of the "I" clone of cDNA,

50 Figure 4 shows a comparison between the nucleotide sequences of the "I" and "N2" clones,

Figure 5 shows a comparison between the amino-acids of the "N2" gene and  $\Delta$ 12 desaturases of *Arabidopsis* and of soya,

55 Figure 6 shows the homology between hazel Δ12 desaturase and various desaturases of other plants both plastid and of the endoplasmic reticulum,

Figure 7 shows the expression of the N2 gene in various varieties of hazel both in the leaves and in the kernels.

# Isolation and cloning of the FAD2 gene of Arabidopsis thaliana for use as a probe

In order to isolate the gene which codes for hazel  $\Delta 12$  desaturase enzyme, it was necessary to use the FAD2 gene of *Arabidopsis* as a probe.

In order to isolate the Arabidopsis gene, two oligonucleotides were used as "primers" for the amplification of the sequences included between the start and the end of the gene. The oligonucleotides used were NOCC1 (CTGAATTC-CAGGTGGAAGAATGCC) which contains the Eco RI restriction site and the sequences corresponding to the portion between bases 100 and 116 of the gene (Okuley J. et al, 1994, The Plant Cell 6, 147-158) and NOCC4 (AGGAATTC-GACAATTTCTTCACCATCATGC) which contains the restriction site of the Eco RI enzyme and the sequences complementary to the portion between base 1245 and base 1266. The amplification reaction was as follows: 12.8µl H<sub>2</sub>O, 2.5µl 10 x PCR buffer (Perkin Elmer), 2.5µl Arabidopsis genome DNA(10 ng/l), 1µl dNTP, each 2.5mM, 2µl 25mM MgCl<sub>2</sub>, 1µl NOCC1 oligonucleotide (50ng/μl), 1μl NOCC4 oligonucleotide (50ng/μl) 0.2μl Taq I DNA polymerase (Perkin Elmer) (5U/µl). The mixture thus prepared was subjected to 1 denaturing cycle for 1 minute at 94°C and to 40 cycles composed as follows: 30 seconds at 94°C, 1 minute at 52°C, 2 minutes at 72°C. The amplification products were separated on 1% agarose gel in TAE buffer (0.04M Tris-acetate, 0.002M EDTA) and stained with ethidium bromide at a concentration of 0.5μg/ml. The portion of gel containing the fragment of the expected length was withdrawn. In order to extract the DNA, 10μl of Qiaex resin (Qiaex extraction kit, firm Qiagen) were added for each 200mg of gel. The supplier's method was then followed. The DNA was then supplemented with a tenth of a volume of 10XH buffer (Boehringer) and 20 units of Eco RI enzyme (Boehringer). After incubation overnight at 37°C, the DNA was precipitated with 0.1 volumes of 5M NH<sub>4</sub>OAc and one volume of isopropanol. After 10 minutes at ambient temperature, the DNA was centrifuged for 20 minutes at 14000 rpm and the precipitate was washed with 70% ethanol. The DNA was resuspended in 15µl of H<sub>2</sub>O. The concentration was determined on gel by comparison with a known standard.

The amplified fragment was inserted in the pUC18 vector. A ligation mixture was prepared as follows: 1µl pUC18 plasmid DNA cut with Eco RI (20ng), 1.5µl fragment amplified with NOCC1 and 4 (25ng), 1µl 10X ligase buffer (Boehringer), 1µl T4 DNA ligase (1U/µl) (Boehringer), 4.5µl H<sub>2</sub>O. The reaction mixture was incubated at 14°C for 12 hours.

In order to prepare competent cells, the method based on the compound hexamino-cobalt chloride was used (Maniatis, Molecular cloning, 1989, Cold Spring Harbor Laboratory Press, 1.76-1.81). 10μl of the ligation mixture were added to each aliquot of competent cells, defrosted on ice. After the cells had been incubated on ice for 30 minutes they were subjected to thermal shock at 42°C for 90 seconds and were then replaced in ice for 60 seconds. After the addition of 0.5 ml of SOC broth (2% Bactotryptone, 0.5% yeast extract, 10mM NaCl, 2.5mM KCl, 10mM MgCl<sub>2</sub>; 20mM glucose, pH7), the cells were incubated at 37°C with stirring for 90'. 100, 200 and 300 μl aliquots were spread on plates containing solid LB broth (10gr/l NaCl, 10gr/l Bactotryptone, 5gr/l yeast extract, pH7.5, 15gr/l agar) with the addition of 50μg/ml of ampicillin and in the presence of IPTG and X-Gal. The plates were then incubated at 37°C overnight.

Some of the bacterial colonies obtained were first analyzed for their plasmid content by a quick method (Maniatis, Molecular cloning, 1989, Cold Spring Harbor Laboratory Press, 1.32). The colonies containing a plasmid of the expected length were grown and their plasmid DNA extracted (Maniatis, Molecular cloning, 1989, Cold Spring Harbor Laboratory Press, 1.33). Those containing a fragment of the expected length (1160 bp) were identified by digestion of the plasmid DNA with Eco RI. The E1 colony was selected.

One end of the insert of the E1 colony was sequenced. The plasmid DNA of the E1 clone was denatured and partially sequenced by Sanger's method using the enzyme Sequenase and  $^{35}$ S-dATP (Amersham). The sequencing products were separated on 8% acrylamide, 8M urea, 1XTBE gel. After electrophoresis, the gel was dried and exposed overnight in contact with an autoradiographic plate ( $\beta$  max, Amersham). The sequence was compared with that published and was identical, identifying the *Arabidopsis* FAD2 gene in the cloned fragment.

#### s Extraction of nucleic acids from hazel

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Hazelnuts of the Nocchione, Montebello and San Giovanni varieties were harvested when almost fully ripe. The kernel was skinned before being used or frozen in liquid nitrogen. The leaves were harvested at a young stage and frozen in liquid nitrogen. 3 ml of extraction buffer were used for each gram of vegetable material with the use of the method described by Verwoerd et al. (Nucl. Ac. Res., 1989, 2362). Upon completion of the extraction, two selective precipitations were carried out by the addition of NaCl 2M, and 2 volumes of 95% ethanol to eliminate polysaccharides. The final pellet was resuspended in  $H_2O$ . Further centrifuging was then carried out to eliminate any non-resuspended material.

On the other hand, DNA was extracted from young leaves of the Nocchione and Montebello varieties. The vegetable tissue was pulverized in liquid nitrogen and the DNA extracted by the CTAB (REF) method. To eliminate the polysaccharides, NaCl 2M and 2 volumes of 95% ethanol were added. The samples were incubated for 15' at -80°C and centrifuged for 15' at 4°C and 14000 rmp (Eppendorf). This selective precipitation was repeated twice and the final pellet was resuspended in  $H_2O$ . Further centrifuging was then carried out to eliminate any non-resuspended material.

#### Checking of the probe on hazel DNA and RNA

About 20 $\mu$ g of DNA of the Montebello and Nocchione varieties was cut with Eco RI restriction enzyme in a volume of 300  $\mu$ l in the presence of 400 units of enzyme and H buffer (Promega), with incubation for one night at 37°C. After digestion had been checked by gel electrophoresis of one twentieth of the reaction mixture, the samples were precipitated with ethanol and resuspended in 30 $\mu$ l of H<sub>2</sub>O. The DNA was then subjected to electrophoresis on 0.7% agarose gel and transferred by capillarity onto nylon membrane (Southern blot) for one night in the presence of 20 x SSC (3M NaCl, 0.3M Na citrate). The membrane was dried in air for 30' and then fixed by UV treatment (120,000  $\mu$ J/cm²).

The *Arabidopsis*  $\Delta$ 12 desaturase gene was used as a probe. For this purpose, the plasmid DNA of the E1 clone (5µg) was cut with 20 units of Eco RI in the presence of H buffer (Boehringer) in a volume of 30µl for 12 hours at 37°C. The insert of the clone was separated from the vector by electrophoresis on 1% agarose gel and extracted from the gel with the use of Qiaex resin in accordance with the suppliers' instructions (Qiagen). The DNA was denatured for 10' at 100°C, cooled rapidly in dry ice, and marked by the random priming method with the use of 6000 Ci/mmol ( $\alpha$ <sup>32</sup>)P dATP and the reagents of Boehringer's marking kit.

The nylon membrane containing the hazel DNA was prehybridized for 1.5 hours at 55°C in standard buffer (5 x SSC, 0.1% (w/v) N-laurylsarcosine, 0.02% SDS, 1% blocking reagent solution) (10% blocking reagent solution: 10gr Boehringer blocking reagent in 150mM NaCl, 100mM maleic acid, pH7.5). The membrane was then hybridized with the *Arabidopsis* probe for one night at 55°C. The non-hybridized probe was washed twice for 15' in 2 x SSC, 0.1% SDS and twice for 15' each in 0.3 x SSC, 0.1% SDS, always at a temperature of 55°C. The probe remained coupled to the homologous sequences on the membrane was detected by autoradiography.

The RNA extracted from the young leaves of the Montebello and Nocchione varieties and from the kernels of the San Giovanni variety was separated on denaturing gel in the presence of formamide and transferred to nylon membrane by Northern blotting (Maniatis, Molecular cloning, 1989, Cold Spring Harbor Laboratory Press, 7.43-7.45).  $40\mu g$ /sample of total RNA extracted from San Giovanni kernels, Nocchione leaves and Montebello leaves were used. 60 pg of probe were used as a positive control. The RNA was loaded onto a 1% agarose gel in the presence of formal-dehyde. The samples were then subjected to electrophoresis for 3 hours at 80 volts in the presence of 1xMOPS. The gel was rinsed in  $H_2O$  and then stained with ethidium bromide  $0.5 \mu g$ /ml to display the RNA. The RNA was then transferred onto a nylon membrane (Boehringer) by "capillary blotting" in the presence of  $20 \times SSC$  throughout the night at  $4^{\circ}C$ . After transfer, the membrane was dried on 3 MM paper and then fixed by crosslinking using UV light (Stratagene UV Stratalinker  $120000 \mu J/cm^2$ ). The RNA was hybridised with the Arabidopsis  $\Delta 12$  desaturase probe as described for the DNA. Detection was carried out by autoradiography. The heterologous Arabidopsis probe was able to display a band with a molecular weight of about 1500 bp in the hazel RNA and 3 bands of about 18, 8 and 2.8 kb in the hazel DNA cut with Eco RI.

#### Construction of a gene library of cDNA

The gene library of cDNA was constructed from RNA from kernels harvested when almost fully ripe and taken from plants of the San Giovanni variety. For this purpose, the Poly(A)+mRNA was isolated from the total RNA with the use of the Poly(A)Tract mRNA Isolation System II, in accordance with the method provided by the firm Promega. The samples were eluted in  $\rm H_2O$  and precipitated with 0.1 volumes of 3M NaOAc and 3 volumes of 95% ethanol. After one night at -80°C, the RNA was centrifuged for 15' at 14000 rpm (Eppendorf), the pellet was rinsed in 75% ethanol and resuspended in  $\rm 10\mu l$  of  $\rm H_2O$ . The concentration was read with a spectrophotometer and the yield was 3.2 $\rm \mu g$  of Poly(A)+mRNA per mg of total RNA.

The messenger RNA polyadenilate derived from kernels of the San Giovanni variety was used as a template for the synthesis of complementary DNA (cDNA) with the use of Boehringer's "cDNA synthesis kit" in accordance with the method recommended by the suppliers. An extraction was then carried out with one volume of phenol:chloroform: isoamyl alcohol (25:24:1). The cDNA was then purified in a Pharmacia column (cDNA spun columns) after the addition of NaCl 100 mM. The buffer used was the following: 10mM Tris-HCl pH 7.5, 1mM EDTA, 150mM NaCl. Eco RI "adaptors" (Pharmacia) were added to the ends of the cDNA. The reaction mixture contained: 5µl of cDNA (half of the cDNA obtained from 6μg of Poly(A)+RNA), 10μl of ligase buffer 10 x (Promega), 10μl of Eco RI adaptors (0.01μ/μl), 6 units of T4 DNA ligase (Promega), in a final volume of 100µl. After incubation for 12 hours at 12°C, the ligase enzyme was inactivated for 10' at 65°C. Phosphorylation of the adaptors then followed by the addition, to the 100µl mixture, of 10µl of 100mM ATP and 10 units of T4 polynucleotide kinase. After incubation at 37°C for 30', the enzyme was inactivated by incubation for 10' at 65°C. Purification was then carried out with one volume of phenol:chloroform:isoamyl alcohol (25:24:1). The cDNA was then purified from fragments of less than 400 bp as follows. After the addition of NaCl to a final concentration of 0.1M NaCl, the cDNA was separated by chromatography in a column with Sepharose CL-4B resin (Size prep 400 spun column, Pharmacia) according to the method suggested by the suppliers. The fragments of cDNA shorter than 400 bases were thus excluded. The cDNA was precipitated with one thirtieth of a volume of 3M NaOAc and 2 volumes of 95% ethanol, centrifuged and resuspended in 10µl of H2O.

The cDNA was inserted in the λ phage vector Zap II cut with Eco RI and dephosphorylated (Stratagene) in the following manner: 2μl of cDNA (200 ng), 1μl of λ Zap II cut with Eco RI (1μg/μl) (Stratagene), 0.5μl of T4 DNA Ligase (4U/μl) (Promega), 0.5μl of 10 x ligation buffer (Promega), 1μl of H<sub>2</sub>O. The reaction mixture was incubated for 14 hours at 12°C. The mixture containing the cDNA inserted in the vector was used for the reconstruction of the phages with the use of Stratagene's Gigapack Gold "in vitro packaging" kit. The gene library of phages thus obtained was constituted by about 300,000 pfu (plaque-forming units). In order to amplify the gene library, XL1 Blue MRF' cells were prepared as described by Stratagene and used the same day. The gene libraries were plated at a concentration of about 5000 pfu per plate (95 cm²). After growth, the phages were resuspended in SM (5.8gr/l NaCl, 2gr/l MgSO<sub>4</sub>.7H<sub>2</sub>O, 50ml/l 1M Tris HCl (pH 7.5), 5ml/l 2% gelatine) and, after the addition of chloroform to 5% and incubation for 15 minutes at ambient temperature, the cell debris was centrifuged for 10 minutes at 2000 x g. Chloroform to 0.3% was added to the supernatural liquid and the phages were preserved at 4°C. Aliquots were preserved at -70°C after the addition of DMSO to 7%. The gene library was titled.

#### Construction of a partial genome gene library

The DNA of the Nocchione variety was digested with Eco RI restriction enzyme and separated on agarose gel. The fragments with lengths of up to 10000 bp (base pairs) were isolated from the gel with the use of Qiaex resin according to the Qiagen's method. For cloning in the  $\lambda$  vector Zap II, 400ng of DNA fragments were incubated with 1 $\mu$ g of desphosphorylated  $\lambda$  Zap II (Stratagene) in the presence of ligase buffer and 1.5 units of T4 DNA ligase (Promega) for 12 hours at 14°C.

Strategene's Gigapack Gold "in vitro packaging" kit was used in accordance with the suppliers' instructions to make up the gene library. The gene library of phages thus produced was amplified as described for the cDNA gene library. The complexity of the gene library was 1,500,000 clones. This gene library was also amplified.

#### Screening of the cDNA gene library

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About 250,000 phages of the cDNA gene library were plated on LB broth in the presence of XL1 Blue MRF' cells, divided into 12 plates each containing 20,000 pfu. After growth, the phages were transferred onto nylon membranes and their denatured DNA was fixed on the membranes as described by Boehringer for screening with non-radioactive probes. The membranes were then hybridized with the *Arabidopsis* Δ12 desaturase gene. The probe was prepared by the isolation of the insert containing the entire coding region of the gene from the plasmid. The insert was then marked with digoxigenin-dUTP with the use of Boehringer's "DNA labelling kit". Prehybridization was carried out in standard buffer (Boehringer) and hybridization was carried out in the same buffer with the addition of the *Arabidopsis* probe at a concentration of 10ng/ml and at a temperature of 55°C.

After washing twice in 2xSSC, 0.1% SDS for 5 minutes at ambient temperature and washing twice in 0.3xSSC, 0.1%SDS at 55°C, detection was carried out with the use of an anti-digoxigenin antibody conjugated with alkaline phosphatase (Boehringer) and a chemiluminescent substrate (AMPPD, Boehringer).

11 positive phage plaques were identified. These were isolated, the phages resuspended in SM and titled. From 50 to 200 phages were plated for each positive plaque. The plaques were transferred onto nylon membranes and subjected to a second hybridization with the *Arabidopsis*  $\Delta$ 12 desaturase probe, as already described above. The following clones which could hybridize with the *Acabidopsis*  $\Delta$ 12 desaturase gene were obtained from the second screening: I, F  $\Delta$ 

#### Screening of the genome gene library

The gene library of Nocchione DNA was subjected to screening in the same way as the cDNA gene library. 1,600,000 phages were plated, divided into 40 plates. After growth, they were transferred to nylon membranes as described for the cDNA gene library. The membranes were then hybridized with the *Arabidopsis* Δ12 desaturase gene as described for the cDNA gene library. Autoradiography of the membranes showed 9 positive plaques. These plaques were isolated, titled and subjected to a second screening. 6 plaques were re-confirmed as positive. 4 of these gave a very strong signal.

# Analysis of the clones isolated

The following positive phage clones were converted into plasmids by *in vivo* excision in accordance with the method suggested by Stratagene (Gigapack Gold in vitro packaging): I, F, 4 (cDNA gene library), N2, N11, N17, N18, N21, N25 (genome gene library).

The plasmid DNA of the clones of the cDNA gene library was isolated and the length of the insert analyzed by digestion with Eco RI. The plasmid DNA of the genome clones was isolated, the length of the insert analyzed by cutting

with restriction enzyme, and the clones rechecked by hybridization with the *Arabidopsis* probe. Figure 1 shows the map of the N2 genome clone.

#### Sequencing

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The N2 clone was selected from the genome clones. For sequencing, the insert was fragmented with Sau3A restriction enzyme and the fragments obtained were subcloned in pUC18 vector cut with BamHI (Maniatis, Molecular cloning, 1989, Cold Spring Harbor Laboratory Press, 1.68-1.69). The clones obtained were analyzed both for the length of the insert and by hybridization with the *Arabidopsis* probe. Since the N2 insert was 2.8 kb and hence longer than the Δ12 desaturase gene, the hybridization excluded the clones containing sequences outside the gene. The insert of the I, F, 4 and N2 clones was isolated and sequenced with the use of the Sequenase kit and (35S)dATP. All of the clones (cDNA and genome) were first sequenced at the ends with the use of primers which could couple with the vector in both orientations. In order to complete the internal regions and to assemble the fragments of the N2 genome clone, internal oligonucleotides were then designed and synthesized and were used for the sequencing. The following table shows the sequences of the internal oligonucleotides:

OLIGONUCLEOTIDE	SEQUENCE
N2-3SS	CAG ACC AGC ATC CGA GAC
N2-3SD	GGA TTG GCT TAG GGG GGC
N2-29R'S	GCC AAC CAT GTC ATC AAC CC
NOCCS	ATG GTA GAG AAG AGA TGG TG
COL	CTG GTG GGT TGT TGA AG
N2-S1N	GGA GAG GTC ATA AAC AAC

The I and F clones were sequenced entirely. As far as the N2 clone is concerned, only the regions corresponding to the gene were sequenced. Figures 2 and 3 show their sequence. The I and F cDNA clones were identical. A comparison between I and the N2 genome clone showed the same sequence (Fig. 4), indicating that N2 contains the gene which codes for the cDNA of the I clone.

#### Comparison between the gene isolated and other desaturases

The nucleotide and amino-acid sequence of the N2 clone was compared with other desaturases (Figure 6). The greatest homology was with the two  $\Delta 12$  desaturases of the endoplasmic reticulum and with a hydroxylase of ricin which uses the same substrate as  $\Delta 12$  desaturase. Homology with the plastid  $\Delta 12$  desaturases and with both the plastid and endoplasmic reticulum  $\Delta 15$  desaturases was, however, much lower. Figure 5 shows the comparison between the amino-acid sequence of hazel  $\Delta 12$  and those of Arabidopsis and soya.

#### Checking of the expression of the hazel $\Delta 12$ desaturase gene

RNA was extracted from kernels of the San Giovanni, Montebello and Nocchione varieties and from leaves of the Montebello and Nocchione varieties. After separation on agarose gel, the RNA was transferred onto a nylon membrane and hybridized with the insert of the I clone marked with digoxigenin. The result is shown in Figure 7, in which a band is visible in the kernel RNA but not in that of the leaves.

# SEQUENCE LISTING

5	(1) GENERAL INFORMATION:
3	<ul> <li>(i) APPLICANT:</li> <li>(A) NAME: SOREMARTEC S.A.</li> <li>(B) STREET: Dreve de l'Arc-en-Ciel 102</li> <li>(C) CITY: Arlon-Schoppach</li> </ul>
10	(E) COUNTRY: Belgium (F) POSTAL CODE (ZIP): 6700
	(ii) TITLE OF INVENTION: Isolation and sequencing of the hazel FAD2-N gene
15	
	(iii) NUMBER OF SEQUENCES: 4
20	<pre>(iv) COMPUTER READABLE FORM:     (A) MEDIUM TYPE: Floppy disk     (B) COMPUTER: IBM PC compatible     (C) OPERATING SYSTEM: PC-DOS/MS-DOS     (D) SOFTWARE: PatentIn Release #1.0, Version #1.30</pre>
	(EPO)
25	<pre>(vi) PRIOR APPLICATION DATA:    (A) APPLICATION NUMBER: CH 0550/96    (B) FILING DATE: 04-MAR-1996</pre>
30	(2) INFORMATION FOR SEQ ID NO: 1:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 1662 base pairs (B) TYPE: nucleic acid
	(C) STRANDEDNESS: double
35	(D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: DNA (genomic)
	· (iii) HYPOTHETICAL: NO
40	(iv) ANTI-SENSE: NO
	(vi) ORIGINAL SOURCE:
	<ul><li>(A) ORGANISM: Corylus avellana cv. Nocchione</li><li>(F) TISSUE TYPE: leaves</li></ul>
45	(vii) IMMEDIATE SOURCE: (B) CLONE: N2
50	<pre>(ix) FEATURE:</pre>

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		(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	: 1:					
5		ATAA 60	IA AA	AGTA	AGCTO	TTA	TACC	TCA I	AGTA	GGGT"	rt cc	TAT	'GACA	TAA 1	GAGT	ccc
		TCCT 20	TT T	TATO	GAGGT	r GCT	'ATAA	TTG (	CAAA'	TGTC	CA AA	TCAT	AGGC	ATA	TGGA	TCC
10		ACTA 80	TT A	ATAT	PATG:	r agi	GTGT	TTT'	тттт	TTTC	сс тс	TAAA	ATT	C TCT	CACA	ССТ
	AAGT	TGAT 23		CTCC	AGCA	тт	GACA	ATAGO	СТС	CTGT	AGAC			GA G Ly Al		
15													1			
		CGA 281														
	Ser	Arg	Met	Pro	Ala	Thr	Asn	Lys	Pro	Lys	Glu	Gln	Lys	Thr	Pro	He
20	5					10					15					20
		CGA 329	)													
25	Gln	Arg	Ala	Pro	His	Thr	Lys	Pro	Pro	Phe	Thr	Leu	Ser	Gln	Leu	Lys
25					25					30					35	5
<i>30</i>		GCC 377	7													
30	Lys	Ala	Val	Pro	Pro	Asn	Сув	Phe	Gln	Arg	Ser	Leu	Leu	Arg	Ser	Phe
				40					45	<b>i</b>				5	0	
35		TAT 42	5													
	Ser	Tyr	Val	Val	Tyr	Asp	Leu	Ser	Leu	Ala	Phe	Leu	Phe	Tyr	Tyr	Ile
			55					60	)				6	5		
40													•			
	GCT	ACC		TAC	TTC	CAT	CTC	CTC	CCI	CAC	ccc	CTT	TCC	TAC	TTG	GCA
	Ala	47 Thr	3 Ser	Tyr	Phe	His	Leu	Leu	Pro	His	Pro	Leu	Ser	Туг	Let	Ala
45		70					79	5				8	0			
		TCA 52	1													
50	Trp	Ser	Ile	е Туг	Tr	Ala	a Leu	ı Glr	Gly	у Суя	: Ile	Lev	ı Thi	r Gly	y Val	L Trp
	85	5				90	)				95	<b>i</b>				100

	GTC	ATC		CAT	GAG	TGC	GGT	CAC	CAT	GCC	TTT	AGT	GAC	TAC	CAA	TGG
	Val	569 Ile		His	Glu	Cys	Gly	His	His	Ala	Phe	Ser	Asp	Tyr	Gln	Trp
5					105					110					115	
		GAT 617														
10	Val	Asp	Asp	Met	Val	Gly	Leu	Thr	Leu	His	Ser	Ala	Leu	Leu	Val	Pro
				120		•			125					130		
15	TAC	TTT 665		TGG	AAG	ATT	AGC	CAC	TGT	CGC	CAC	CAC	TCT	AAC	ACC	GGC
	Tyr	Phe		Trp	Lys	Ile	Ser	His	Cys	Arg	His	His	Ser	Asn	Thr	Gly
			135					140					145	<b>.</b>		•
20	TCC	CTT		CGA	GAT	GAG	GTG	TTT	GTC	ссс	AAG	CCG	AAA	TCC	AAA	ATG
	Ser	713 Leu		Arg	Asp	Glu	Val	Phe	Val	Pro	Lys	Pro	Lys	Ser	Lys	Met
25		150					155	•				160	)		•	
		TGG 761	L													
	Pro	Trp	Phe	Ser	Lys	Tyr	Phe	Asn	Asn	Pro	Pro	Gly	Arg	Val	Leu	Thr
30	165					170					175				٠	180
	CTT	TTG 809		ACA	CTC	ACT	CTA	GGC	TGG	ccc	TTG	TAC	TTA	GCC	TTG	AAT
35	Leu	Leu		Thr	Leu	Thr	Leu	Gly	Trp	Pro	Leu	Tyr	Leu	Ala	Leu	Asn
					185					190	)				19	5
40	GTT	TCT 85		CGA	CCC	TAT	GAT	CGT	TTT	GCT	TGC	CAC	TAŢ	GAT	CCC	TAT
	Val	Ser		Arg	Pro	Tyr	Asp	Arg	Phe	Ala	Суѕ	His	Tyr	Asp	Pro	Tyr
				200					205	5				21	0	
45		CCC 90	5													
	Gly	Pro	Ile	Tyr	Ser	Asn	Arg	Glu	Arg	Cys	Gln	Ile	Phe	· Val	Ser	Yab
50			215	1				220	)				22	5		
	GCT	GGT 95		TTI	GCI	ACA	A ACI	TAT	GTG	CTI	TAC	TAC	C GCP	A GCA	ATG	TCA

	Ala	Gly	Val	Phe	Ala	Thr	Thr	Tyr	Val	Leu	Tyr	Tyr	Ala	Ala	Met	Ser
		230					235					240				
5				<b>a</b>										mmc	C.M.C	2002
		1001						TTC								
	Lys	Gly	Leu	Ala	Trp	Leu	Val	Phe	Ile	Tyr	Gly	Met	Pro	Leu	Leu	
10	245					250					255					260
	GTG	AAT 1049		TTC	CTT	GTA	TTA	ATC	ACC	TAC	TTG	CAG	CAC	ACT	CAC	CCT
15	Val	Asn	Gly	Phe	Leu	Val	Leu	Ile	Thr	Tyr	Leu	Gln	His	Thr	His	Pro
					265					270	)				275	5
	GCA			CAC	TAT	GAC	TCA	TCA	GAA	TGG	GAT	TGG	CTT	AGG	GGG	GCA
20	Ala	109 Leu	7 Pro	His	Tyr	Asp	Ser	Ser	Glu	Trp	Asp	Trp	Leu	Arg	Gly	Ala
				280					285	i				290	)	
25		114	5					TAC								
	Leu	Ala	Thr	Ala	Asp	Arg	Asp	Tyr	Gly	Met	Leu	Asn	Lys	Val	Phe	His
30			295					300	)				30	5		
	TAA	ATC 119		GAC	ACC	CAT	GTG	GCT	CAC	CAT	CTC	TTC	TCT	ACC	ATG	CCT
	Asn	Ile	Ile	. Asp	Thr	His	Val	. Ala	His	His	Leu	Phe	Ser	Thr	Met	Pro
35		310	•				31	5				32	0			
	CAT	TAC		r GC	OTA A	GAA	GCC	ACC	AA.	A GCA	ATC	AAG	TCA	ATA	TTC	GGC
40	His	Tyr	Hie	s Ala	a Met	: Glu	a Ala	a Thr	. Lys	s Ala	a Ile	Lye	Ser	Ile	Lev	Gly
	325	5				330	)				335	5				340
45	AAA			C CA	G TT	r GAT	r GG	C ACT	r cc	A GT	TAC	C AAC	G GCI	A GTO	TG	AGG
40	Lys	128 Ty:	39 c <b>Ty</b> :	r Gl	n Phe	e Ası	p Gl	y Thi	Pr	o Va	1 Туі	c Ly	s Ala	a Val	Tr	Arg
					34	5				35	0				3	55
50		17	37													r AAC
	G1	u Al	a Ly	s Gl	u Cy	s Le	и Ту	r Va	1 G1	u Se	r As	p Gl	u Gl	y Al	a Pr	o Asn
<i>EE</i>																

		360		365	.3*	70
5	1390		AT CAG AGC A	AAG CTG TGA TA	ATTGGCTGG A	TAGAGCCAA
	33	75	380	)		
10	AGAAAATGTO	G ATTAGTAAG	G TAGTGTCTTT	GGTCAGTTTG G	TGTGTTAAG G	ААСАААТАА
15	TAATAATTA( 1510	G CGACTATGA	A TAGTTATTGI	тааасаааат т	CACCCTTAT G	TTTAGCAGG
	AACTTTTCT( 1570	G GCTACACTT	T TTTTCGTATO	AAAAGCGCAT A	Т ТААТТТТТТ	GTTATATTG
20	TTTTGACAT	T ACTCAAGCT	т сааааттаат	T ATCACAGAAA A	TATCCAATG T	'CGAAGGTTT
	CATTGTAGG 1662		TT ATATTGAGO	GT GG		
25	(2) INFOR	MATION FOR	SEQ ID NO:	2:		
30	(i	(A) LENGTI (B) TYPE:	CHARACTERIS  1: 383 amino amino acid  OGY: linear	STICS: o acids		
	(ii) (xi)	MOLECULE T	YPE: protein ESCRIPTION:	SEQ ID NO: 2	:	
35	Met Gly A 1	la Arg Ser 5	Arg Met Pro	Ala Thr Asn 10	Lys Pro Ly	s Glu Gln 15
40	Lys Thr P	ro Ile Gln 20	Arg Ala Pro	His Thr Lys 25	Pro Pro Ph	e Thr Leu 30
	Ser Gln L	eu Lys Lys 35		Pro Asn Cys	Phe Gln Ar 45	g Ser Leu
45	Leu Arg S 50	Ser Phe Ser	Tyr Val Val	l Tyr Asp Leu	Ser Leu Al 60	a Phe Leu
50	Phe Tyr 1	Tyr Ile Ala	Thr Ser Ty:	r Phe His Leu 75	Leu Pro Hi	s Pro Leu. 80
	Ser Tyr 1	Leu Ala Trp 85		r Trp Ala Leu 90	Gln Gly Cy	ys Ile Leu 95

	Thr	Gly	Val	Trp	Val	Ile	Ala	His	Glu 105		Gly :	His	His	Ala F 110	he S	er
5	Asp	Tyr	Gln 115	Trp	Val .	Asp	Asp	Met 120		Gly	Leu	Thr	Leu 12!	His S	Ser 1	Ala
10	Leu	Leu 130	Val	Pro	Tyr	Phe	Ser 135		Lys	Ile	Ser	His 140	Cys )	Arg I	His 1	His
	Ser 145	Asn	Thr	Gly	Ser	Leu 150	Asp	Arg	qaA	Glu	Val 155	Phe	Val	Pro 1	Lys :	Pro 160
15	Lys	Ser	Lys	Met	Pro 165	Trp	Phe	Ser	Lys	Tyr 170	Phe )	Asn	Asn	Pro	Pro 17	Gly 5
20	Arg	Val	Leu	Thr 180	Leu	Leu	Ile	Thr	Leu 189	Thr 5	Leu	Gly	Trp	Pro 19	Leu 0	Tyr
	Leu	Ala	Leu 195		Val	Ser	Gly	Arg 200	Pro )	туг	Asp	Arg	Phe 20	Ala 5	Cys	His
25	Tyr	Asp 210		Туг	Gly	Pro	11e 21	Tyr 5	Ser	Asn	Arg	Glu 22	Arg 0	Cys	Gln	Ile
30	Phe 225		Ser	Asp	Ala	Gly 230	Val	Phe	Ala	Thr	Thr 235	Tyr	Val	Leu	Tyr	Tyr 240
	Ala	Ala	a Met	Ser	Lys 245		Leu	Ala	Trp	Leu 25	Val 0	Phe	Ile	Tyr	Gly 25	Met 5
35	Pro	Let	ı Lev	1   Ile 260		Авг	ı Gly	Phe	Leu 26	Val	Leu	Ile	Thr	Туг 27	Leu '0	Gln
40	Hie	s Thi	r His		Ala	Le:	ı Pro	His 28	5 Туі 10	as T	Ser	Ser	Glu 2	Trp 85	Asp	Trp
	Le	u Ar		y Ala	a Lev	ı Ala	a Thi 29	r Ala 95	а Абј	o Ar	g Asp	Ту1 3	Gly 00	/ Met	Leu	Asn
45	Ly 30		l Ph	e Hi	s Ası	n Il 31	e Il	e As	p Th	r Hi	s Va:	l Ala	a Hi	s His	Leu	Phe 320
	Se	r Th	r Me	t Pr	o Hi: 32		r Hi	s Al	a Me	t Gl 3	u Ala 30	a Th	r Ly	s Ala	Ile 3	Lys 35
50	Se	er Il	e Le	u Gl	y Ly	s Ту	r Ty	r Gl	n Ph	e As	p Gl	y Th	r Pr	o Val	L <b>Ту</b> і	Lys

5	Ala Val Trp Arg Glu Ala Lys Glu Cys Leu Tyr Val Glu Ser Asp Glu 355 360 365
10	Gly Ala Pro Asn Lys Gly Val Phe Trp Tyr Gln Ser Lys Leu * 370 375 380
	(2) INFORMATION FOR SEQ ID NO: 3:
15	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 1133 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>
·	(ii) MOLECULE TYPE: cDNA to mRNA
20	(iii) HYPOTHETICAL: NO
	(iv) ANTI-SENSE: NO
•	(v) FRAGMENT TYPE: C-terminal
25	<ul><li>(vi) ORIGINAL SOURCE:</li><li>(A) ORGANISM: Corylus avellana L. cv. San Giovanni</li><li>(D) DEVELOPMENTAL STAGE: Seed, storage deposition stage</li></ul>
30	(Vii) IMMEDIATE SOURCE: (B) CLONE: I
35	<pre>(ix) FEATURE:     (A) NAME/KEY: mRNA     (B) LOCATION:11133     (D) OTHER INFORMATION:/partial     /gene= "Fad2"</pre>
33	(ix) FEATURE:
40	(A) NAME/KEY: CDS (B) LOCATION:11019 (D) OTHER INFORMATION:/partial /codon_start= 3 /product= "delta-12 desaturase" /gene= "Fad2"
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:
	TC CAA CGC TCT CTC CTA CGC TCG TTC TCA TAT GTT GTT TAT GAC CTC
	Gln Arg Ser Leu Leu Arg Ser Phe Ser Tyr Val Val Tyr Asp Leu
50	385 390 395
	TCC TTA GCC TTC CTC TTC TAC TAT ATT GCT ACC TCT TAC TTC CAT CTC

		95	5													
	Ser		Ala	Phe	Leu	Phe	Tyr	Tyr	Ile	Ala	Thr	Ser	Tyr	Phe	His	Leu
5		400					405					410				
									001	<b></b>				<b></b>	0.0m	ama.
		143														
10	Leu	Pro	His	Pro	Leu	Ser	Tyr	Leu	Ala	Trp	Ser	He	Tyr	Trp	Ala	
	415					420					425					430
	CAA		TGC	TTA	CTC	ACC	GGC	GTT	TGG	GTC	ATC	GCA	CAT	GAG	TGC	GGT
15	Gln	19: Gly	Cys	Ile	Leu	Thr	Gly	Val	Trp	Val	Ile	Ala	His	Glu	Сув	Gly
					435					440					445	5
20		23														
	His	His	Ala	Phe	Ser	Asp	Tyr	Gln	Trp	Val	Asp	Asp	Met	Val	Gly	Leu
				450					455					460	)	
25	ACC	CTT 28	CAC	TCT	GCT	CTT	TTA	GTT	CCA	TAC	TTT	TCA	TGG	AAG	ATT	AGC
	Thr		His	Ser	Ala	Leu	Leu	Val	Pro	Tyr	Phe	Ser	Trp	Lys	Ile	Ser
30			465					470					475	5		
	CAC	mcm	CGC	CAC	CAC	mcm	አእሮ	N.C.C	ccc	መርር	CTTT	GAC	CCA	CAT	GAG	GTG
		33	5													Val
35	ure	-	_	нів	nis	ser			GIY	261	Den			veb	GIU	Val
		480					485	•				49	U			
	TTT	GTC 38		AAG	CCG	AAA	тсс	AAA	ATG	CCA	TGG	TTT	TCT	AAG	TAC	TTC
40	Phe			Lys	Pro	Lys	Ser	Lys	Met	Pro	Trp	Phe	Ser	Lys	Tyr	Phe
	495					500	)				505					510
												3 mc		cmc	<b>3</b> C/F	C C C C
45		43	31													CTA
	Asn	Asr	) Pro	Pro	Gly	Arç	y Val	Leu	Thr			Ile	Thr	Leu	Thr	Leu
					515	5				52	0				52	.5
50		41	79													GAT
	Gly	Tr	Pro	Leu	тул	Let	ı Ala	Lev	Ası	ı Val	Ser	G13	y Arg	Pro	Туг	Asp

				530					535			:		540		
5	CGT	TTT 527		TGC	CAC	TAT	GAT	ccc	TAT	GGC	ccc	ATT	TAT	TCC	ТАА	CGC
	Arg	Phe		Сув	His	Tyr	Asp	Pro	Tyr	Gly	Pro	Ile	Tyr	Ser	Asn	Arg
			545					550					555			
10															-	
		AGG 575														
	Glu	Arg	Суѕ	Gln	Ile	Phe	Val	Ser	Asp	Ala	Gly	Val	Phe	Ala	Thr	Thr
15		560					565					570				
	TAT	GTG 623		TAC	TAC	GCA	GCA	ATG	TCA	AAA	GGG	CTG	GCA	TGG	CTT	GTA
	Tyr	Val		Tyr	Tyr	Ala	Ala	Met	Ser	Lys	Gly	Leu	Ala	Trp	Leu	Val
	575					580		-			585			•		590
•	TTC	ATT 671		GGT	ATG	CCA	TTG	CTC	ATA	GTG	AAT	GGC	TTC	CTT	GTA	TTA
25	Phe	Ile	Tyr	Gly	Met	Pro	Leu	Leu	Ile	Val	Asn	Gly	Phe	Leu	Val	Leu
					595					600					60	5
30		719	9													TCA
	Ile	Thr	Tyr	Leu	Gln	His	Thr	His	Pro	Ala	Leu	Pro	His	Tyr	Asp	Ser
				610					615	•				620	D	
35	TCA	GAA 76		GAT	TGG	CTT	AGG	GGG	GCA	ТТG	GCG	ACG	GCG	GAT	AGA	GAT
	Ser	Glu	Trp	Asp	Trp	Leu	Arg	Gly	Ala	Leu	Ala	Thr	Ala	Asp	Arg	Asp
40			625	•				630	)				63	5		
	TAC			CTG	AAT	AAG	GTT	TTC	CAC	TAA	ATC	ATA	GAC	ACC	CAT	GTG
	Туг	81 Gly		. Leu	Asn	Lys	Val	Phe	His	Asn	Ile	Ile	Asp	Thr	His	Val
45		640					645	5				65	0			
	GCI	CAC		r CTC	TTC	TCI	ACC	: ATG	CCI	CAT	TAC	CAT	GCA	ATO	GAA	GCC
50	Ala	His	His	s Lev	ı Phe	Ser	Thr	Met	Pro	His	Туг	His	ala Ala	Met	Gli	ı Ala
	655	5				660	)				665	i i				670

	ACC	AAA 91		ATC	AAG	TCA	ATA	TTG	GGC	AAA	TAC	TAC	CAG	TTT	GAT	GGC
	Thr			Ile	Lys	Ser	Ile	Leu	Gly	Lys	Tyr	Tyr	Gln	Phe	Asp	Gly
5					675					680					685	5
										0.0	000		CNC	mc c	Cmm	mam
		95	9											TGC		
10	Thr	Pro	Val	Tyr	Lys	Ala	Val	Trp			Ala	Lys	Glu	Cys		Tyr
				690					695					700	)	
15		100	7											TGG		
	Val	Glu	Ser	Asp	Glu	Gly	Ala	Pro	Asn	Lys	Gly	Val	Phe	Trp	Tyr	Gln
			705					710	)				71	5		
20	AGC	ממ	: ሮሞር	בטים:	רבי	ייינוכר	TGG	ATAC	AGC	CAA J	AGAAJ	AATG'	TG A'	TTAG	raag:	G
		10	59 Leu													
25				GGTC	AGTT'	TG GI	GTGT	DAAT	GAA	CAAA	TAA 1	TAAT	LTTA.	AG CG	АСТА	TGAA
30	TAG	-	TTGT 133	TAAA												
	(2)	INI	ORM	MOITA	FOF	SEÇ	O I D	NO:	4:							
35			(	SEQU (A) I (B) I	LENGT	TH: 3	339 á ino á	mino acid	STICS o ac	3: ids						
		(i (x	i) Mo i) Si	DLEC! EQUE!	ULE '	TYPE DESCI	: pro	otei:	SEQ	ID :	NO:	4:				
40		n Ar	g Se	r Lei		a Arç	g Ser	Phe	. Ser		Val	Val	Tyr	Asp	Leu	Ser 15
45	Le	u Al	a Ph		u Phe O	е Туг	туг	: Ile	Ala 2	a Thr !5	Ser	Туг	Phe	His	Leu 30	Leu
50	Pr	o Hi		o Le	u Se	r Ty	r Lei		a Try 10	Sei	: Ile	е Туг	Trp	Ala 45	Leu	Gln
	G1	.у Су	s Il	e Le	u Th	r Gl	y Va	1 Tr	p Va	1 11	e Ala	a His	s Glu	ı Cys	: Gly	' His
55																

		50					55						0			
5	His 65	Ala	Phe	Ser	Asp	Tyr 70	Gln	Trp	Val	Asp	Asp 75	: Met	Val	Gly	Leu	Thr 80
	Leu	His	Ser	Ala	Leu 85	Leu	Val	Pro	туг	Phe 90		Trp	Lys	Ile		His 5
10	Cys	Arg	His	His 100	Ser	Asn	Thr	Gly	Ser 105		Asp	Arg	Asp	Glu 11		Phe
15	Val	Pro	Lys 115	Pro	Lys	Ser	Lys	Met 120		Trp	Phe	Ser	Lys 12	Tyr 5	Phe	Asn
20	Asn	Pro 130	Pro	Gly	Arg	Val	Leu 135		Leu	Leu	Ile	Thr 14	_	Thr	Leu	Gly
	Trp 145	Pro	Leu	Tyr	Leu	Ala 150	Leu	Asn	Val	Ser	Gly 155	Arg	Pro	туr	Asp	Arg 160
25	Phe	Ala	Cys	His	Туг 165	Asp	Pro	Tyr	Gly	Pro 170		Tyr	Ser	naA	Arg 17	
	Arg	Cys	Gln	Ile 180	Phe	Val	Ser	Asp	Ala 185	_	Val	Phe	Ala	Thr 19	_	туr
30	Val	Leu	Туг 195	Tyr	Ala	Ala	Met	Ser 200		Gly	Leu	Ala	Trp 20	Leu 5	Val	Phe
35	Ile	Tyr 210		Met	Pro	Leu	Leu 219		Val	Asn	Gly	Phe 22		Val	Leu	Ile
40	Thr 225	Tyr	Leu	Gln	His	Thr 230	His	Pro	Ala	Leu	Pro 235	His	Tyr	Asp	Ser	Ser 240
40	Glu	Trp	qaA	Trp	Leu 245		Gly	Ala	Leu	Ala 25		Ala	Asp	Arg		Tyr 55
45	Gly	Met	Leu		-		Phe			_	Ile	Asp	Thr		Val 70	Ala
50	His	His	Leu 275		Ser	Thr	Met	Pro 28		Tyr	His	Ala		Glu 85	Ala	Thṛ
	Lys	Ala 290		Lys	Ser	Ile	Leu 29		Lys	Туг	Tyr		Phe	Asp	Gly	Thr

Pro Val Tyr Lys Ala Val Trp Arg Glu Ala Lys Glu Cys Leu Tyr Val 305 310 315 320

Glu Ser Asp Glu Gly Ala Pro Asn Lys Gly Val Phe Trp Tyr Gln Ser 325 330 335

Lys Leu \*

10

25

40

5

#### 15 Claims

- 1. A fragment of DNA from hazel (Corylus avellana L.) comprising the nucleotide sequence shown in Figure 2.
- A DNA fragment comprising the nucleotide sequence shown in Figure 2 from base 222 to base 1367, which codes for the hazel Δ12 desaturase enzyme of the endoplasmic reticulum or for a homologous sequence which can code for the same amino-acid sequence.
  - 3. A nucleotide sequence coding for a protein or peptide having an amino-acid homology greater than or equal to 80% and preferably greater than 90% with the hazel △12 desaturase enzyme of the endoplasmic reticulum of Claim 2 and having the function of the said enzyme.
  - 4. A recombinant DNA sequence comprising a DNA sequence according to Claims 1, 2 and 3, or a portion of such a sequence, together with sequences regulating expression.
- 5. A recombinant DNA molecule comprising a cloning vector in which a DNA sequence according to any one of Claims 1, 2, 3 and 4 is inserted.
  - 6. A DNA molecule according to Claim 5, in which the cloning vector is a plasmid or a phage.
- A DNA molecule according to Claim 4 or Claim 5 having the restriction map shown in Figure 1.
  - 8. A host organism including a recombinant DNA molecule according to any one of Claims 3 to 6.
  - 9. A host organism according to Claim 8, selected from a vegetable cell, an animal cell, and a micro-organism.
  - 10. A genetically modified organism capable of expressing the FAD2-N gene, having the amino-acid sequence shown in Figure 2 from bp 222 to bp 1367, portions of this gene, or this gene conjugated with other molecules and containing sequences which can inactivate endogenous genes.
- 45 11. A hazel Δ12 desaturase enzyme of the endoplasmic reticulum having the amino-acid sequence shown in Figure 2 in substantially pure form.
  - 12. A fusion polypeptide comprising the amino-acid sequence of the enzyme of Claim 11, in which the amino-acids additively connected thereto do not interfere with the desaturase activity or can easily be eliminated.
  - 13. The use of the FAD2-N gene coding for the hazel Δ12 desaturase enzyme of the endoplasmic reticulum or of portions thereof for the isolation of enzymes having the function of hazel desaturase or of the desaturase of another species.
- 14. The use of the nucleotide sequences of the FAD2-N gene shown in Figure 2 for the construction of expression systems which can alter the fatty-acid content in hazel.

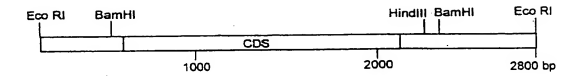


Fig. 1 - Restriction map of the genomic clone "N2". CDS: coding region; bp: base pair.

CCICATAAAAAAGTAAGCTCATTIACCTCAAGTAGGGTTTCCTTATGACAAATGAGTCCC 60 GGAGTATTTTTCATTCGAGTAAATGAGTTCATCCCAAAGGAATACTGTTTACTCAGGG  GCAATCCTTTTCTATGAGGTGCTATAATTGCAAATGTCCAAATCATAGGGATATGATCC 120 CGTTAGGAAAAGATACTCCACGATATTAACGTTTACAGGTTTAGTATCCCTATACCTAGG  AAATACTATTAATATTATGTAGTGTGTTTTTTTTTT		2 - Nucleotide sequence of the gene FAD2-N corresponding n internal fragment of the genomic clone "N2". Aminoacid dues of the coding region are also reported.	to ar
CGTTAGGAAAAGATACTCCACGATATTAACGTTTACAGGTTTAGTATCCCTATACCTAGG  AAATACTATTAATATTATGTAGTGTGTTTTTTTTTT	60	TCATAAAAAAGTAAGCTCATTTACCTCAAGTAGGGTTTCCTTATGACAAATGAGTCCC 6 AGTATTTTTTCATTCGAGTAAATGGAGTTCATCCCAAAGGAATACTGTTTACTCAGGG	C C G G
AAGTTGATATTTTCTCCAGCATTGGACATAGAAAAAAAAA	120	AATCCTTTTCTATGAGGTGCTATAATTGCAAATGTCCAAATCATAGGGATATGGATCC 1	GC CG
TICAACTAAAAGAGGTCGTAACCTGTATCGGAGACATCTGTTACCCTCGATCTTCGGCTT  Met Gly Ald Arg Ser Arg  TGCCTGCTACCAACAAGCCTAAAGAGCAAAAAACACCCATCCAGCGAGCACCACACACA	160	NATACTATTAATATTATGTAGTGTGTTTTTTTTTTTCCCTCAAATTTACTCTCACACCT 1 ITATGATAATTATAATACATCACACAAAAAAAAAAAGGGAGTTTAAATGAGAGTGTGGA	A A T T
Met Pro Ald The Ash Lys Pro Lys Giu Gin Lys The Pro IIe Gin Arg Ald Pro His The  AACCCCATTCACTCTTAGCCAACTCAAGAAAGCCGTCCCACCCA	240	TCAAC TAAAAGAGGTCG TAACCTG TATCGGAGACATC TGTTACCCTCGATC TTCGGCTT	A A T T
Lys Pro Pro Phe Thr Leu Ser Gin Leu Lys Lys Ald Val Pro Pro Ash Cys Pre Gin Arg  CTCTCCTACGCTCGTTCTCATATGTTTGTTTATGACCTCCCTTAGCCTTCCTCTTCTACT 420  GAGAGGATGCGAGCAAGAGTATACAACAAATACTGGAGAGGAATCGGAAGGAGAAGATGA  Ser Leu Leu Arg Ser Phe Ser Tyr Val Vai Tyr Asp Leu Ser Leu Ald Pre Leu Pre Tyr  ATATTGCTACCTCTTACTTCCATCTCCCTCCCTCACCCCCTTTCCTACTTGGCATGGTCAA 480  TATAACGATGGAGAATGAAGGTAGAGGAGGAGGGAGGAGAAGGATGAACCGTACCAGTT  Tyr IIe Ald Thr Ser Tyr Phe His Leu Leu Pro His Pro Leu Ser Tyr Leu Ald Trp Ser  TCTATTGGGCTCTCCAAGGCTGCATTCTCACCGGCGTTTTGGGTCATCGCACATGAGTGCG  GIE Tyr Trp Ald Leu Gin Giy Cys IIe Leu Thr Giy Vai Trp Val IIe Ald His Giu Cys  GTCACCATGCCTTTAGTGACTACCAATGGGTTGATGACATGGTTGGCCTAACCCTTCACT 600  CAGTGGTACGGAAATCACTGTACCAATGGGTTGATGACAACCGGATTGGGAAGTGA  GIY His His Ald Phe Ser Asp Tyr Gin Trp Val Asp Asp Met Val Giy Leu Thr Leu His  CTGCTCTTTTAGTTCCATACTTTTCATGGAAGATTAGCCACTGTCGCACCACTCTAACA 660  GACGAGAAAATCAAGGTTATGAAAAAGTACCTTCTAATCGGTGACAGCCGTGGTGAGAATTGT	300	CGGACGATGGTTGTTCGGATTTCTCGTTTTTTGTGGGTAGGTCGCTCGTGGTGTGTGT	AC
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CGGATGCTGGTGTCTTTGCTACAACTTATGTGCTTTACTACGCAGCAATGTCAAAAGGGC GCCTACGACCACAGAAACGATGTTGAATACACGAAATGATGCGTCGT.TACAGTTTTCCCG	960
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Leu Alc Trp Leu Val Fine ile Tyr Giy Met Pro Leu Leu ile Val Asn Giy Fine Leu Vai	
TAATCACCTACTTGCAGCACACTCACCCTGCATTGCCGCACTATGACTCATCAGAATGGGATTAGTGGATGGGATGAGTGGGACGTAACGGCGTGATACTGAGTAGTCTTACCC	1080
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ATTGGCTTAGGGGGGCATTGGCGACGGCGGATAGAGATTACGGAATGCTGAATAAGGTTT TAACCGAATCCCCCCGTAACCGCTGCCGCCTATCTCTAATGCCTTACGACTTATTCCAAA	1140
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TCCACAATATCATAGACACCCATGTGGCTCACCATCTCTCTC	1200
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Fig. 3 - Nucleotide sequence of cCNA clone "I".

Fig 4 - Nucleotide sequence alignment of clones "I" (I.SEQ) and

"N2" (N2.SEQ). CCTCATAAAAAAGTAAGCTCATTTACCTCAAGTAGGGTTT M2.SEQ CCTTATGACAAATGAGTCCCGCAATCCTTTTCTATGAGGT N2.SEQ GCTATAATTGCAAATGTCCAAATCATAGGGATATGGATCC N2.SEQ 121 AAATACTATTAATATTATGTAGTGTTTTTTTTTTTTCCC NZ.SEQ 161 TCAAATTTACTCTCACACCTAAGTTGATTTTCTCCAGCAT NZ.SEQ 201 TGGACATAGCCTCTGTAGACAATGGGAGCTAGAAGCCGAA N2.SEQ 241 IGCCTGCTACCAACAAGCCTAAAGAGCAAAAAAAACACCCAT N2.5EQ .......... 281 CCAGCGAGCACCACACAAAACCCCCATTCACTCTTAGC N2.5EQ 321 CAACTCAAGAAAGCCGTCCCACCCAATTGTTTCCAACGCT N2.52Q 10 CTCTCCTACGCTCGTTCTCATATGTTGTTTATGACCTCTC I.SEQ 361 CTCTCCTACGCTCGTTCTCATATGTTGTTTATGACCTCTC NO.SEQ 3G CTTAGCCTTCCTCCTACTATATTGCTACCTCTTACTTC I.SEQ 401 CTTAGCCTTCCTCTTCTACTATATTGCTACCTCTTACTTC NZ.SEQ 90 CATCTCCTCCCTCACCCCCTTTCCTACTTGGCATGGTCAA I.SEQ 441 CATCTCCTCCCTCACCCCCTTTCCTACTTGGCATGGTCAA NZ.SEQ 130 TOTATTGGGCTCTCCAAGGCTGCATTCTCACCGGCGTTTG\_I.SEQ 481 TOTATTGGGCTCTCCAAGGCTGCATTCTCACCGGCGTTTG NZ.SEQ 170 GGTCATCGCACATGAGTGCGGTCACCATGCCTTTAGTGAC [.SEQ 521 GGTCATCGCACATGAGTGCGGTCACCATGCCTTTAGTGAC NZ.SEQ ZIO TACCANTGGGTTGATGACATGGTTGGCCTARCCCTTCACT I.SEQ 561 TACCARTGGGTTGATGACATGGTTGGCCTAACCCTTCACT N2.SEQ 250 CTGCTCTTTTAGTTCCATACTTTTCATGGAAGATTAGCCA I.SEQ 601 CTGCTCTTTAGTTCCATACTTTTCATGGAAGATTAGCCA N2.52Q 290 CTGTCCCCACCACTCTAACACCGGCTCCCTTGACCGAGAT I.SEQ 541 CTGTCGCCACCACTCTAACACCGGCTCCCTTGACCGAGAT N2.SEQ 330 GAGGTGTTTGTCCCCAAGCCGAAATCCAAAATGCCATGGT I.SEQ 681 GAGGTGTTTGTCCCCAAGCCGAAATCCAAAATGCCATGGT, N2.52Q 370 TTTCTAAGTACTTCAACAACCCACCAGGTAGGGTCCTCAC 1.SEQ 721 TTTCTAAGTACTTCAACAACCCACCAGGTAGGGTCCTCAC N2.5EQ 410 TCTTTTGATCACACTCACTCTAGGCTGGCCCTTGTACTTA I.SEQ 761 TCTTTTGATCACACTCACTCTAGGCTGGCCCTTGTACTTA N2.SEQ 450 GCCTTGAATGTTTCTGGCCGACCCTATGATCGTTTTGCTT I.SEQ 801 GCCTTGAATGTTTCTGGCCGACCCTATGATCGTTTTGCTT N2.5E0

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 241 RYAAAQGMASMI'CLYGVPLLIVNAFLVLITYLQHTHPSLP L26295.PRC
 280 HYDSSEWDWLRGALATADRDYGMLNKVFHNIIDTHVAHHL N2.PRO
 Z81 HYTSSEWDWLRGALATVORDYGILNKVFHNITOTHVAHHL 143921.7RC
 291 HYDSSEWDWLRGALATVDRDYGILNKVFHNITDTHVAHHL L26296.PRC
 320 FSTMPHYHAMEATKAIKSILGKYYQFDGTPVYKAVWREAK N2.PRO
 321 FSTMPHYHAMEATKAIKPILGEYYRFOETPF<u>VKAMW</u>REAR L43921.PRO
 321 FSTMPHYNAMEATKAIKPILGOYYQFDGTPWYVAMYREAK L26295.FRO
 360 ECLYVES, DEGAPNKGVFWYQSKL
 361 ECIYVEPO'QSTESKGVFWYNNKL
                                                             L43921.PRC -
                                                             L26296.PRO
 361 ECIYVEPDREGDKKGVYWYNNKL
```

Fig. 5 - Aminoacid sequence alignment of  $\Delta 12$  desaturase from hazelnut (N2.PRO), Arabidopsis (L26296.PRO) and soybean (L43921.PRO). Homologous residues are boxed.

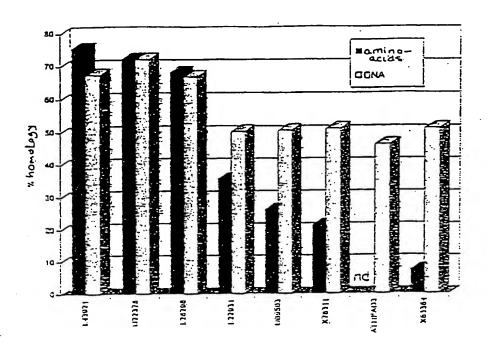


Fig. 5 - Homology between bazel △12 desaturase and other desaturases

143921: Al2 desaturase of the endoplasmic raticulum of soya

U22378: All hydroxylase of ricin L26296: All desaturase of the endoplasmic reticulum of

Arabidopsis theilana
L22931: Als plastid desaturase of Arabidopsis thaliana
U09503: Als plastid desaturase of Arabidopsis thaliana
X78311: Als plastid desaturase of spinach
ATHFAD3: Als desaturase of the endoplasmic reticulum of
Arabidopsis thaliana
X53364: As plastid desaturase of years

X53364: A9 plastid desaturase of rape

Note: nd: not determined since the amino-acid sequence is not known.

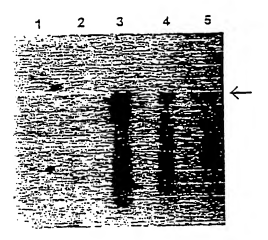


Fig. 7 - Northern blot of RNA of Montabello leaves (line 1), Nocchione leaves (line 2), Montabello kernels (line 3), Nocchione kernels (line 4), and San Giovanni kernels (line 5). The RNA was hybridized with the I clone of cONA.



# **EUROPEAN SEARCH REPORT**

Application Number EP 97 10 3098

1	DOCUMENTS CONSID			
Category	Citation of document with ind of relevant pass	lication, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CL6)
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